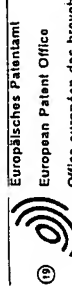


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## BACKGROUND OF THE INVENTION

This invention is directed to a balloon for a blood vessel-dilating catheter.

A balloon catheter having an inflatable balloon secured at its distal end has been applied for various cavities in a living body including a blood vessel. Utility of the balloon catheter is increasing in various medical fields. Of the balloon catheters mentioned above, a blood vessel-dilating catheter is employed in percutaneous transluminal coronary angioplasty (PTCA) to dilate a stenosis or a narrowing in a blood vessel such as coronary artery. In PTCA, femoral artery is secured for example, by Seldinger method; a guiding catheter is introduced into the thus secured femoral artery and advanced through the lumen of the artery until it reaches near the target lesion; the narrowing in the artery, by manipulating a guide wire; a blood vessel-dilating catheter is introduced into the lumen of the guiding catheter to locate the balloon beyond the distal end of the blood vessel-dilating catheter; and a blood vessel-dilating fluid is introduced into the lumen of the blood vessel-dilating catheter to inflate the balloon to thereby dilate the narrowing in the blood vessel.

Such blood vessel-dilating catheter is required to have a trackability so that the blood vessel-dilating catheter can smoothly advance through the lumen of the guiding catheter along the tortuous blood vessel to reach the lesion site. The balloon is required to have a sufficient dimensional stability as well as excellent strength and flexibility so as to avoid excessive dilation of the narrowing of the blood vessel.

Typical balloons for balloon catheters are disclosed in USP 4,093,484; 4,151,244; 4,254,774; 4,906,244; and 5,108,415; and PCT Application No. JP8300202.

The balloons described in these patents and patent applications comprise a mixture of an ethylene-butene-styrene block copolymer and a low molecular weight polyisobutylene having polypropylene optionally added thereto; a composition similar to the one just mentioned wherein butadiene or isoprene is used instead of the ethylene and the butylene; a polyester copolymer; a thermoplastic rubber; a silicone-polycarbonate copolymer; an ethylene-vinyl acetate copolymer; biaxially oriented Nylon 12; biaxially oriented polyethylene terephthalate; polyethylene; a crosslinked ethylene-vinyl acetate copolymer, etc.

The materials particularly used for the balloons of the blood vessel-dilating catheters include polyvinyl chloride (hereinafter abbreviated as PVC), polyethylene (hereinafter abbreviated as PE), biaxially oriented Nylon 12 (hereinafter abbreviated as N12), and biaxially oriented polyethylene terephthalate (hereinafter abbreviated as PET).

Among these, aliphatic high polymers such as PE, PVC, and N12 are highly flexible, realizing a sufficient trackability. These materials, however, are insufficient in their strength to detract from dimensional stability. PET, on the other hand, has excellent strength and dimensional stability. PET, however, has an excessively high modulus of elasticity due to crystallization caused by the biaxial orientation, and therefore, is inferior in impact strength, tear resistance and flexibility, leading to poor trackability of the catheter.

Furthermore, PET is poor in coating adaptability, adhesiveness, and heat sealability to suffer from insufficient properties and workability in preparing the balloon catheter. In addition, PET inherently lacks antithrombotic properties, and it would be quite difficult to subject the PET to various treatments to impart biocompatibility, in particular, blood compatibility.

## SUMMARY OF THE INVENTION

The present invention has been achieved in view of the above-described situation. An object of the present invention is to provide a balloon for a blood vessel-dilating catheter, wherein the softness and the flexibility is improved without compromising the dimensional stability. Another object of the present invention is to provide a balloon for a blood vessel-dilating catheter wherein the modulus of elasticity is reduced to prevent an injury of the blood vessel for a prolonged period of time.

According to the present invention, there is provided a balloon for a blood vessel-dilating catheter fabricated from a biaxially oriented film of an aromatic polyamide or an alloy thereof, said balloon having a calculated modulus of elasticity of from 70 to 190 kg/mm<sup>2</sup>. The aromatic polyamide may preferably be a polyamide prepared by polycondensing xylylenediamine with the aromatic dicarboxylic acid, more preferably adipic acid.

The aromatic polyamide may preferably be a polyamide prepared by polycondensing isophthalic acid and an aliphatic diamine.

The aliphatic diamine may preferably be hexamethylenediamine. The aliphatic dicarboxylic acid may preferably be adipic acid.

The aromatic polyamide may preferably be a polyamide prepared by polycondensing isophthalic acid and an aliphatic diamine.

The aliphatic diamine may contain up to 50% by weight of an aliphatic polyamide.

The aliphatic polyamide blended in the polyamide alloy may preferably be at least a member selected from

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the group consisting of Nylon 6, Nylon 64, Nylon 65, Nylon 66, Nylon 610, Nylon 612, Nylon 9, Nylon 11, Nylon 12, and polyethylene terephthalate.

The balloon may have a burst pressure of at least 10 kg/cm<sup>2</sup>.

#### BRIEF DESCRIPTION OF THE DRAWINGS

Fig. 1 is a sectional view of a blood vessel-dilating catheter having the balloon of the present invention at its distal end.

Fig. 2 is a sectional view taken on line I-I of Fig. 1.

#### DETAILED DESCRIPTION OF THE INVENTION

10 PET as described above, has a high strength, a high modulus of elasticity and a good dimensional stability, although it is poor in flexibility. PET also has a quite high melting point rendering the balloon molding difficult.

On the other hand, aliphatic polymers such as PE, PVC and NY2 are highly flexible, and the resulting blood vessel-dilating catheter may have a good tractability. Such aliphatic polymers, however, are poor in strength and dimensional stability, and also, suffer from decrease in their strength and dimensional alteration upon water impregnation.

In contrast, the balloon for a blood vessel-dilating catheter (hereinafter simply referred to as balloon) of an 20 present invention comprises an aromatic polyamide, which has an excellent dimensional stability, or an alloy thereof with an aliphatic polyamide, which has an excellent flexibility and elasticity. A balloon having both excellent dimensional stability and high flexibility is thereby provided. One example of the balloon of the present invention is shown in Fig. 1, the blood vessel-dilating catheter having the balloon of the present invention at its

distal end consists of an inner tube 1, an outer tube 2 and a balloon 3.

The inner tube 1 has a first lumen 4 with an opening provided at its distal end. The first lumen 4 is intended 25 to transmit a guide wire therethrough.

The outer tube 2 is intended to transmit the inner tube 1 therethrough and has its distal end provided back from the distal end of the inner tube. The inner surface of the outer tube 2 and the outer surface of the inner tube 1 form a second lumen 6. The distal end of the second lumen 6 is connected with the proximal end of the balloon 3 which will be described later, and filled with a fluid (angiographic agent, for example) for inflating the balloon 3. The distal end of the outer tube 2 is fixed to a liner tube 1 without blocking up the second lumen 6. More illustratively, as shown in Fig. 2, it is fixed by a filler 5 provided between the outer tube 2 and the inner tube 1, and the filler 5 has a partial cavity 5a, with which the second lumen 6 and the distal end of the balloon 3 are connected with each other.

The balloon 3 is foldable, and can be folded around the inner tube 1 when it is not inflated. The balloon 3 has a substantially cylindrical portion 3a having almost the same diameter and at least partially cylindrical so 30 that it can dilate a narrowing in a blood vessel with ease. The proximal end 8 of the balloon 3 is fixed liquid-tight to the distal end of the outer tube 2, and the distal end 7 thereof is fixed liquid-tight to the distal end of the inner tube 1 so that a dilating space 15 is formed between the inner surface of the balloon 3 and the outer surface of the inner tube 1. The proximal end of the dilating space 15 is connected with the second lumen 6 through the cavity 5a of the filler 5.

A reinforcing material 14, provided on the outer surface of the inner tube 1 is made of coil spring, and is located near the distal end of the outer tube 2 and also near the center of the balloon 3 on the outer surface 35 of the inner tube 1 so that the position of the balloon 3 can be observed through an X-raying.

Such a balloon having excellent dimensional stability as well as high flexibility may be introduced into the blood vessel with little impact against the blood vessel inner surface to prevent the blood vessel inner surface from being injured.

The reduced modulus of elasticity results in a highly flexible, soft balloon which may be easily folded to a 40 small size with no rigid fold being formed by folding. Hard materials like PET having a high modulus of elasticity are difficult to fold into a small size due to rigid folds formed upon folding.

The balloon catheter having the balloon of the present invention secured thereto has a good tractability to follow the tortuous blood vessel and enables the balloon to reach the target lesion. More illustratively, the tractability of the balloon catheter depends not only the foldability of the balloon to a small size but also on the flexibility of the folded balloon, namely, shell. The balloon formed of PET is poor in the flexibility of the shell to result in an inferior tractability of the catheter. In contrast, the shell of the balloon of the present invention is soft and flexible to realize a good tractability of the catheter.

The balloon of the present invention has a good adhesion to the catheter body due to the properties in-

herent to the resin material. Such a good adhesion to the catheter body is quite favorable for production, and the resulting good adhesion strength between the balloon and the catheter body prevents the balloon from being peeled off from the catheter body during its storage or use.

Furthermore, the balloon of the present invention is excellent in blood compatibility, namely, antithrombotic properties due to the properties inherent to the resin material, and therefore, may be employed within the blood vessel for a prolonged period compared to conventional balloon catheters. In addition, since the balloon of the present invention has good compatibility with other resins as well as good coating adaptability, the balloon may be surface treated with various agents and resin coatings to realize sustained effects of the treatment.

For example, the balloon may have its exterior surface treated with various antithrombotic materials and agents to impart the balloon with high blood compatibility for a prolonged period. The surface of the balloon may be subjected to other surface treatments for other purposes including smooth passage of the blood vessel interior through the lumen of the blood vessel filled with viscous blood, and prevention of the blood vessel interior surface from being injured by the traffic of the balloon.

Although the above-mentioned aliphatic polymers such as PE, PVC and NY2 are flexible, they have low calculated modulus of elasticity and low burst pressure. On the other hand, the balloon made from the above-mentioned PET has a high burst pressure. However, once the PET balloon undergo bursting, it disrupts into numerous small pieces or debris which are quite difficult to recover. In contrast, the balloon of the present invention has a high burst pressure and would not burst even when a pressure of about 10 atmosphere is applied to expansion of the balloon, and even when the balloon should burst, it tears in a wadding configuration to enable a safe recovery.

Also, the balloon of the present invention has an impact strength higher than that of the PET balloon, and therefore, may fully endure a rapid inflation leading to safety.

The balloon for blood vessel-dilating catheter of the present composition comprises either an aromatic polyamide having a good dimensional stability, or an alloy of such an aromatic polyamide with other resin components. In particular, an aliphatic polyamide having a sufficient flexibility or elasticity. When the balloon comprises a polyamide alloy, ratio of the components may be adjusted to realize desired properties. Including a calculated modulus of elasticity in the range of from 70 to 150 kg/mm<sup>2</sup>, and preferably from 70 to 160 kg/mm<sup>2</sup>.

The term aromatic polyamide used herein designates a polyamide produced by polycondensing a diamine and a dicarboxylic acid, at least a part of the diamine or the dicarboxylic acid containing an aromatic ring.

The diamine or the dicarboxylic acid molecules containing an aromatic ring may preferably constitute from 25 to 75% by weight, and most preferably from 40 to 50% by weight of the aromatic polyamide. It is to be noted that the molar ratio of the diamine or the dicarboxylic acid molecules containing no aromatic ring may not necessarily be 1:1. It is also to be noted that the aromatic polyamide may be produced by polycondensing two or more types of diamines and dicarboxylic acids.

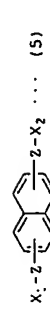
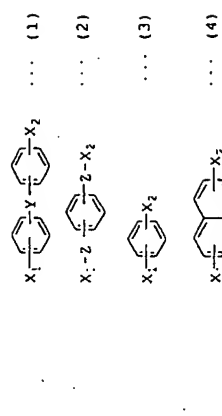
Typical diamines or dicarboxylic acids having at least one aromatic ring include those represented by general formulae [1]:

40

45

50

55



. . . (11).

wherein X<sub>1</sub> and X<sub>2</sub> independently represent -COOH or -NH<sub>2</sub>.  
 Y is a divalent group selected from -O-, -S-, -SO<sub>2</sub>-, -(CH<sub>2</sub>)<sub>n</sub>-O-, -O(CH<sub>2</sub>)<sub>n</sub>-O-.

Z is a divalent group selected from -O(CH<sub>2</sub>)<sub>n</sub>-O-, wherein n is 1 to 4, and -(CH<sub>2</sub>)<sub>n</sub>-.



is benzene ring which may be substituted at any position; and



is naphthalene ring which may be substituted at any position.  
 Among these, those represented by formulae (2) and (3) are preferred.

Typical aliphatic diamines or dicarboxylic acids having no aromatic ring therein include diamines and dicarboxylic acids derived from a straight-chain, a branched, or an alicyclic hydrocarbon, as represented by general formulae [II]:

X<sub>1</sub>-(CH<sub>2</sub>)<sub>n</sub>-X<sub>2</sub>, wherein n is 2 to 12 . . . (1)

X<sub>1</sub>-(CH<sub>2</sub>)<sub>n</sub>-H-(CH<sub>2</sub>)<sub>m</sub>-X<sub>2</sub>, wherein n is 0 to 3 . . . (2)

X<sub>1</sub>-(CH<sub>2</sub>CH<sub>2</sub>O)<sub>m</sub>-X<sub>2</sub>, wherein m is 1 to 1,000 . . . (3)

X<sub>1</sub>-(CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>O)<sub>m</sub>-X<sub>2</sub>, wherein m is 1 to 1,000 . . . (4)

X<sub>1</sub>-(CH<sub>2</sub>CH(CH<sub>3</sub>)O)<sub>m</sub>-X<sub>2</sub>, wherein m is 1 to 1,000 . . . (5)

a dimeric acid . . . (6)

wherein X<sub>1</sub> and X<sub>2</sub> independently represent -COOH or -NH<sub>2</sub>.

55

X<sub>1</sub>- (CH<sub>2</sub>C(CH<sub>3</sub>)<sub>2</sub>CH<sub>2</sub>)<sup>m</sup>-X<sub>2</sub>, wherein m is 1 to 4 . . . (7)

. . . (11).

5 wherein X<sub>1</sub> and X<sub>2</sub> independently represent -COOH or -NH<sub>2</sub>.

Among these, aliphatic diamines or dicarboxylic acids represented by formula (1) are preferred. The aromatic polyamide may have a polymerization degree of approximately 50 to 5,000, and most preferably, approximately 100 to 3,000, and an average molecular weight of approximately 3,000 to 10,000, and most preferably, approximately 5,000 to 20,000.

10 Of the aromatic polyamides mentioned above, the most preferred in view of workability and physical properties are Nylon MXD6 synthesized from m-xylylenediamine and adipic acid and Nylon 61 synthesized from hexamethylenediamine and isophthalic acid.

In the present invention, the aromatic polyamide as described above may be used either alone or as a main component in a polymer alloy wherein the aromatic polyamide is alloyed with a resin component having a sufficient flexibility and elasticity.

Exemplary alloying resins having a sufficient flexibility which may be used alone or in combination of two or more include thermoplastic resins such as aliphatic polyamides, modified polyolefins, polyphenylene oxides, ABS resins and polyesters. Among these, aliphatic polyamides are most preferable in view of their good compatibility with the aromatic polyamide as well as their sufficient workability.

15 Illustrative aliphatic polyamides which may be used alone or in combination of two or more include Nylon 6, Nylon 64, Nylon 66, Nylon 610, Nylon 612, Nylon 46, Nylon 11, Nylon 11, Nylon 12, and polyethylene amide. The term, alloy or polymer alloy used herein is a concept including polymer blend, graft copolymer, random copolymer, block copolymer, and the like.

An alloying agent or a compatibilizing agent may optionally be employed in alloying the aromatic polyamide with other resins such as an aliphatic polyamide.

The resin having a sufficient flexibility, which may typically be an aliphatic polyamide, may comprise up to 50% by weight, most preferably from 0 to 40% by weight of the polymer alloy. When the flexible resin component comprises more than 50% by weight, the resulting balloon will be poor in its modulus of elasticity and strength leading to insufficient dimensional stability.

20 The balloon of the present invention may be secured to the blood vessel-delivering catheter body, which may typically comprise a resin material such as polyvinyl chloride and polyethylene, by thermal fusion using a suitable heating means or with an adhesive or a solvent such as epoxy resin or cyanacrylate adhesive. The balloon of the present invention has an excellent adhesiveness with the catheter body owing to the properties inherent to the resin material, and exhibits excellent adhesion strength after securing the balloon to the catheter body. Use of the balloon of the present invention, therefore, is quite advantageous for the production of a balloon catheter, and the thus produced balloon catheter may be safely stored and used with no risk of the balloon from being peeled off the catheter body.

The balloon of the present invention is produced by biasially orienting the above-described aromatic polyamide or the alloy thereto wherein the aromatic polyamide is the main constituent.

25 In an exemplary process for producing the balloon of the present invention, a tube or a tubular body is fabricated from the above-described aromatic polyamide or the alloy thereof wherein the aromatic polyamide is the main constituent; and the thus produced tube is axially oriented by such means as elongation or drawing. The axial orientation may preferably be carried out at an elevated temperature of, for example, from 45 to 130°C.

30 The thus axially oriented tube may have a length larger than its preorientation length by a factor of about 1.5 to 5.

Next, a mold having a cavity of a configuration corresponding to the balloon in its inflated state is placed over the axially oriented tube at approximately central position in its axial direction. The mold is then heated to a temperature of, for example, from 45 to 130°C to heat the tube. The tube is inflated in its radial direction at the heated portion by applying an elevated pressure. The radius of the tube before the inflation may be about two to eight times larger than the radius of the tube after the inflation.

35 The heated, pressurized conditions of the tube as described above are maintained for a certain period, for example, one second to five minutes, and then, the tube is allowed to cool to approximately room temperature to form the desired balloon configuration. It is to be noted that the balloon may be subjected to repeated cycles of heat application and cooling to thereby remove the strain of the balloon.

After the cooling of the tube, the pressure is reduced to normal pressure, the mold is removed, and the

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balloon is trimmed to produce the balloon of the present invention.

The thus produced balloon of the present invention may have a calculated modulus of elasticity of from 70 to 150 kg/mm<sup>2</sup>, and more preferably from 70 to 160 kg/mm<sup>2</sup>.

A balloon having a calculated modulus of elasticity of less than 70 kg/mm<sup>2</sup> is insufficient in strength and dimensional stability. A balloon having a calculated modulus of elasticity in excess of 150 kg/mm<sup>2</sup> is insufficient in softness and flexibility leading to poor trackability of the catheter. Such a balloon also requires a high pressure for inflation, and even when inflated, the balloon may surpass its yield point and experience a plastic deformation to disperse restoration to its original configuration.

The term, calculated modulus of elasticity, E used herein is determined from a calculated tensile strength, Sc which represent the tensile strength in radial direction in film equation. The calculated tensile strength, Sc may be determined by equation [1]:

$$Sc = P \times D / 2t \quad [1]$$

wherein Sc is the calculated tensile strength of the film,

P is the pressure applied;

D is the initial diameter of the balloon, and

t is the thickness of the balloon.

In practice, the calculated tensile strength, Sc and the calculated modulus of elasticity, E are calculated after measuring the stress (pressure) and the strain (balloon diameter) of a balloon filled with water. Detailed measuring processes are described in Examples.

The calculated modulus of elasticity, E corresponds to the slope of the linear portion, wherein Hooke's law is applicable, of the stress-strain curve obtained by plotting the stress component (strength, Sc) in relation to the strain component (inflation of the balloon). In other words, the calculated modulus of elasticity is initial modulus of elasticity of the balloon, which may be determined by equation [1]:

$$E = 8Sc \times D / 8D$$

wherein E is the calculated modulus of elasticity,

8Sc is increment in the strength,

D is initial diameter of the balloon, and

8D is increment in the balloon diameter.

The balloon of the present invention may have a non-limited thickness, which may preferably be from 5 to 30 µm, and more preferably be from 7 to 20 µm.

Conventional PET balloons, which are provided with excellent strength and dimensional stability, are quite hard, and have a calculated modulus of elasticity of 200 kg/mm<sup>2</sup> or higher. In the present invention, the calculated modulus of elasticity has been reduced to 70 to 190 kg/mm<sup>2</sup>, and preferably, to 70 to 160 kg/mm<sup>2</sup>, by fabricating the balloon from the aromatic polyamide or the alloy thereof, whereby a production of a balloon provided with softness and flexibility as well as dimensional stability is enabled.

It is to be noted that the PET balloon could be impaired with a reduced modulus of elasticity by reducing the degree of orientation. In such a case, however, the stress-strain curve would exhibit a yield point, beyond which the dimensional stability as well as the strength would undergo a significant decrease. A pressurization of the balloon beyond such a yield point would lead to a plastic deformation of the balloon upon which a re-storation to its original configuration and dimension would be impossible or render the withdrawal or recovery of the balloon difficult. Therefore, only a considerably limited range of pressure could actually be employed for the PET balloon inflation.

In contrast, the balloon of the present invention is provided with sufficient softness and flexibility without compromising the dimensional stability and the strength. Accordingly, the inner surface of the blood vessel to which the blood vessel-dilating catheter is inserted is prevented from being injured by the balloon upon such an occasion as insertion of the catheter.

In addition, the balloon of the present invention, which is fabricated from the aromatic polyamide or an alloy thereof, has an excellent blood compatibility or anti-thrombotic property, and therefore, may be indwelled in the blood vessel for a prolonged period.

The balloon of the present invention may preferably have a burst pressure of 10 kg/cm<sup>2</sup> or higher, and more preferably, from 13 to 20 kg/cm<sup>2</sup>. The pressure normally required for inflating the balloon is approximately 7 to 8 atm. The balloon of the present invention, which has a burst pressure of 10 kg/cm<sup>2</sup> or higher, would endure a more severe pressurization, than such a normal pressurization, and therefore, could be successfully employed for treating a tight stenosis requiring even higher pressurization.

The present invention is described by referring to the following non-limiting Examples of the present invention, as well as Comparative Examples.

## EXAMPLES

## Example 1

Nylon MXD6 (grade 8121, manufactured by Mitsubishi Gas Chemical Company, Inc.), which is an aromatic polyamide produced by polycondensing m-xylylenediamine and adipic acid, was molded into tube having an inner diameter of 0.7 mm and an outer diameter of 1.1 mm. The tube was axially oriented to a length three times larger than its original length in an atmospheric cavity at a temperature of 81°C. The tube was then placed in a metal cylinder provided with a cylindrical cavity with an inner diameter of 3 mm having opposite tapered ends. The metal cylinder was heated to a temperature of 85°C, and nitrogen was introduced into the tube to a pressure of 15 kg/cm<sup>2</sup> from its opposite ends. The tube was kept at this pressure and temperature for 15 seconds. The tube was then allowed to cool to room temperature in 1 minute with the pressure being kept at the constant level.

The metal cylinder was heated again with the pressure being kept at the constant level, but this time to a temperature of 130°C, and the tube was allowed to heat for 20 seconds and cool to room temperature in 50 seconds.

After reducing the pressure, the biaxially oriented balloon was removed from the metal cylinder, and trimed to obtain the balloon of the present invention. The resulting balloon had an outer diameter at its dilated portion of 3 mm and a film thickness of 15.0 µm.

## Example 2

With 80% by weight of the Nylon MXD6 employed in Example 1 was blended and kneaded 20% by weight of Nylon 6 (grade 103BFR), manufactured by Unitika Ltd., which is an aliphatic polyamide, in a twin-screw extruder to produce MXD6/N6 alloy pellets.

A balloon was produced from these pellets in a manner similar to Example 1. The resulting balloon had an outer diameter at its dilated portion of 3 mm and a film thickness of 15.5 µm.

## Example 3

With 80% by weight of the Nylon MXD6 employed in Example 1 was blended and kneaded 20% by weight of Nylon 6 (grade 103BFR), manufactured by Unitika Ltd., which is an aliphatic polyamide, in a twin-screw extruder to produce MXD6/N6 alloy pellets.

A balloon was produced from these pellets in a manner similar to Example 1. The resulting balloon had an outer diameter at its dilated portion of 3 mm and a film thickness of 14.3 µm.

## Example 4

With 80% by weight of the Nylon MXD6 employed in Example 1 was blended and kneaded 20% by weight of Nylon 6, which is an aliphatic polyamide, in a twin-screw extruder to produce N6/N6 alloy pellets.

A balloon was produced from these pellets in a manner similar to Example 1. The resulting balloon had an outer diameter at its dilated portion of 3 mm and a film thickness of 15.1 µm.

## Example 5

The procedure of Example 1 was repeated except that Nylon 61 produced by polycondensing hexamethylene diamine and isophthalic acid was used for the aromatic polyamide. The resulting balloon had an outer diameter at its dilated portion of 3 mm and a film thickness of 15.3 µm.

## Comparative Example 1

With 70% by weight of the Nylon 61 employed in Example 4 was blended and kneaded 30% by weight of Nylon 6, which is an aliphatic polyamide, in a twin-screw extruder to produce N6/N6 alloy pellets.

A balloon was produced from these pellets in a manner similar to Example 1. The resulting balloon had an outer diameter at its dilated portion of 3 mm and a film thickness of 15.3 µm.

## Comparative Example 2

A commercially Nydor® II balloon Cordis-Helix™ (manufactured by United States Catheter and Instrument) Having an outer diameter at its dilated portion of 3 mm and a film thickness of 8 µm.

## Comparative Example 2

A commercially Nydor® II balloon Cordis-Helix™ (manufactured by Dardis) having an outer diameter at its dilated portion of 3 mm and a film thickness of 8 µm.



8. The balloon according to any one of claims 1 to 7 wherein said balloon has a burst pressure of at least 10 N/mm<sup>2</sup>.

5 10 15 20 25 30 35 40 45 50 55 60 65 70 75 80 85 90 95 100

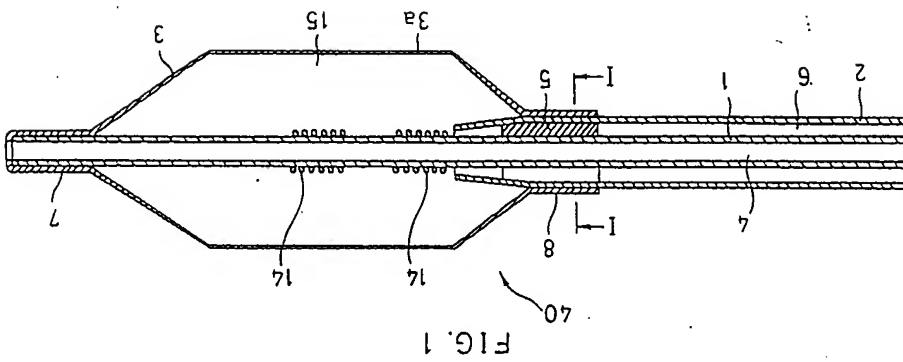


FIG. 1

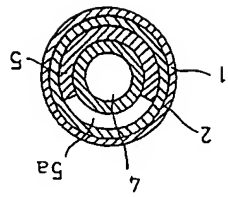


FIG. 2

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EUROPEAN SEARCH REPORT

Application Number

EP 92 40 2739

DOCUMENTS CONSIDERED TO BE RELEVANT		CLASSIFICATION OF THE APPLICATION (if Cls.)
Category	Character of document with indication where appropriate of relevant passages	Relevant to claims 1-7 A61L29/00
Y	EP-A-0 362 826 (CORDIS CORPORATION) * column 13, line 30 - column 15, line 14; claims *	
Y	DATABASE NP1L Week 8128, Derwent Publications Ltd., London, GB; AN 81-508870 & JPA-55 062 129 (TOYOB0 KK) * abstract *	1-7
		TECHNICAL FIELDS SEARCHED (in Cls.)
		A61L
<p>The present search report has been drawn up for all claims.</p> <p>Place of search Date of completion of the search Requester</p> <p>THE HAGUE 15 JANUARY 1993 G. COUSINS-VAN STEEN</p>		
<p>CATEGORY OF CITED DOCUMENTS</p> <p>X : periodically repeated if it has been found Y : found in connection with another document of the same category A : technically interesting O : not directly relevant P : irrelevant</p>		
<p>I : primary or principal document E : embodiment R : reference D : document cited in the application L : document cited for other reasons M : material document H : historical document C : commercial document B : bibliographical reference G : general document P : patent publication T : technical document S : standard document U : unexamined document D : document containing drawings F : foreign language document N : non-patent literature W : work of fiction O : other document</p>		